

BRMICH, Stefan; TABENSKI, Zbigniew

Free skin transplantation in the treatment of large varicose leg
ulcers. Polski przegl. chir. 26 no.9:825-836 Sept 54.

1. w Oddzialow Chirurgicznych Szpitala Miejskiego w Gliwicach.
Ordynatorzy: dr. S.Brmich & dr. Z.Tabenski
(VARICOSE VENES, ulcers
surg. skin grafts)
(SKIN TRANSPLANTATION
free graft in ther. of varicose ulcers)

ERMICH, Stefan; TABENSKI, Zbigniew

Surgical indications in gastric and duodenal ulcers. Polski tygod. lek. 11 no.28:1252-1257 9 July 56.

1. Gliwice Plac "olnosc 8/3.
(PEPTIC ULCER, surgery,
indic. (Pol))

TABENSKI, Zbigniew; ERMICH, Stefan

Comparative evaluation of methods of treating myelitis.
Polski przegl. chir. 28 no.11:1155-1161 Nov 56.

1. Z Oddzialow Chirurgicznych Szpitala Miejskiego w Gliwicach.
I. Gliwice, pl. Wolnosci 8, m. 3.
(MYELITIS, ther.
penicillin & surg. (Pol))
(PENICILLIN, ther. use
myelitis (Pol))

ERMICH, Stefan; ZAWISLAK, Mieczyslaw

Acute inflammation of the tail of the pancreas. Polski przegl.
chir. 33 no.7/9:974-975 '61:

1. Z Oddzialu Chirurgicznego Ogolnego Szpitala Miejskiego w Gliwicach
Ordynator: dr S. Ermich.
(PANCREATITIS)

ERMICH, Stefan; LEJANKA, Maria; KANIEWICZ, Zbigniew

Section of Oddi's sphincter. Pcl. przegl. chir. 35 no.7/8:
806-807 '63.

1. Z Oddzialu Chirurgii Ogolnej Szpitala Miejskiego w
Gliwicach Ordynator: dr S. Ermich.
(VATER'S AMPULLA) (SURGERY, OPERATIVE)

ERMICH, Stefan, dr.; RUTKOWSKI, Boleslaw, dr.

Diagnostic and prognostic value of tongue picture in acute abdominal diseases. Pol. przegl. chir. 37 no.2:103-109 F '65.

J. z Oddzialu Chirurgicznego Ogolnego Szpitala Miejskiego w Gliwicach (Ordynator: dr. S. Ermich) i z Oddzialu Anestezjologicznego Szpitala Miejskiego w Gliwicach (Ordynator: dr. B. Rutkowski).

SHANIN, Yu.N.; BURMISTROV, M.I.; BALYUZEK, F.V.; ERMILOV, N.I.

Surgery on the open heart with the D. Milrose apparatus, Vest.
khir. 84 no.1:129-132 Ja '60. (MIRA 13:10)
(PERFUSION PUMP (HEART))

L 23144-66 EWP(j)/I/EWP(t)/ETC(m)-6 LIP(c) M/W/R
ACC NR: AF6010708 SOURCE CODE: CZ/0034/65/000/001/028R/0289

AUTHOR: Styblo, Karel (Engineer); Ermis, Frantisek; Pivoda, Petr (Graduate chemist); Kovarik, Milos

ORG: VZU NHKG VZKG, Ostrava

TITLE: Determination of gases, and oxygen particularly, by means of the instrument exhalograph EA-1

SOURCE: Hutnicke listy, no. 4, 1965, 288-289

TOPIC TAGS: steel, aluminum, metal chemical analysis, laboratory instrument

ABSTRACT: The instrument is supplied by Balzers of Liechtenstein. Description of the instrument is given. Operation of the apparatus is described. The results are reproducible, and obtained in 3 minutes. In samples of steel stilled with Al (up to 0.05% Al) the time required is 5-6 minutes; when 0.5 Al is present the time needed is 10-12 minutes. At higher Al contents, up to 20 minutes is needed for the analysis. Orig. art. has: 2 figures and 1 table. [JIRIS]

SUB CODE: 11, 07 / SUBM DATE: none / OTH REF: 006

Card 1/1 04R

A Study of Electrolytically Isolated Carbides from Low-Alloy Boiler Plate. M. Bichu, A. Smirnov, and V. Ernits. (Hungarian Lit., 1955, 19, (2), 149-162). [In Czech] - The separation and micro-analysis of the carbides are described. Chemical and electron-diffraction methods were used for the identification. Carbides in vanadium steels were found to stabilize sooner than in molybdenum steels of similar compositions. The mode of carbide stabilization is described on the basis of data obtained in experiments carried out in the range 800-850° C. over periods of 8000-125,000 hr. - p. 7.

28 ② Jan

RADOMIROV, P., prof.; ERMOLAEV, Iv.; KOZAROVA, M.; KHRISTOV, G.; STOIMENOVA, St.;
NEDEVA, D.

Molybdenum as microfertilizer in Bulgaria. Selskostop nauka 2
no. 9:1153-1160

BRUNSH, N. ; KALNINS, A.

Antisepticization of open-air wooden constructions. p. 139.

БИОЛОГИЧЕСКАЯ НАУКА. СЕЛЬСКОМУ ЛЕСНОМУ КОМПЛЕКСУ. (Latvijas PSR
Zinatnu akademija. Biologijas zinatnu nodala) Riga, Latvia, №. 3, 1957.

Monthly list of East European Acquisitions ("FAI"), DC, Vol. 6, No. 8,
August 1959.
Unclassified.

BELYAKOV, G. (Riga); ERMUSH, N. [Ermusa, N.] (Riga); KALNIN'SH, A.
[Kalinins, A.] (Riga)

Possibilities of utilizing pitch-hydrophobized sand. Vestis Latv ak
no.3:85-90 '61. (EEAI 10:9)

1. Akademiya nauk Latviyskoy SSR, Institut lesokhozaystvennykh
problem i khimii drevesiny.

(Concrete) (Sand)

ERMUSH, N. [Ermusa, N.]

New developments in the field of wood protection. Vestis Latv ak no.9:
139-140 '61.

GROMOV, V.S., kand. khim. nauk, otv. red.; DORBURG, G.E., kand. khim. nauk, red.; IYEVIN'SH, I.K.[Ievins, I.], kand. tekhn. nauk, red.; KAL'NINA, V.K.[Kalnina, V.], kand. tekhn. nauk, red.; RUPAYS, Ye.A.[Rupais, E.], kand. khim. nauk, red.; SERGEYEVA, V.N., doktor khim. nauk, red.; ERMUSH, N.A.[Ermus, N.], st. nauchn. sotr., red.; YUKNA, A.D.[Jukna, A.], kand. tekhn. nauk, red.; LEVI, S., red.; SHKLENNIK, Ch., red.

[Chemical processing and preserving of wood] Khimicheskaya pererabotka i zashchita drevesiny. Riga, Izd-vo AN Latv.SSR, 1964. 238 p. (MIRA 10:1)

1. Latvijas Padomju Socialistiskas Republikas Zinatnu Akademija. 2. Institut khimii drevesiny AN Latviyskoy SSR (for Gromov, Sergeyeva, Ermush).

ERMYAN, A. V.

USSR/Medicine - Synthomycin

Jan/Feb 52

"Dermatitis Produced by Synthomycin," A. V. Emyan, Moscow Clinical Hosp of Infectious Diseases.

"Vest Venerol i Dermstol" No 1, p 51.

Synthomycin, a synthetic prep identical with Synthomycin, a synthetic prep identical with the antibiotic chloromycetin, is being used in the treatment of typhus, typhoid, and dysentery. Author describes the reaction to this drug observed in various cases. Predominating reactions after the injection of synthomycin were skin eruption resembling the rash of measles, itching

222T16

of the epidermis, formation of papules over the body and extremities, erythema with symptoms of measles, headache, and general weakness upon an application of synthomycin paste removal of the dressing revealed papules and itching. In the author's experience, these symptoms disappear within 48 hours after discontinuation of the synthomycin treatment.

222T16

"APPROVED FOR RELEASE: Thursday, July 27, 2000

CIA-RDP86-00513R00041222

ERN, O. S.

ERN, O.S.

Practical work in preparing soil mixtures and peat-humus pots.
Biol. v. shkole no.2:87-89 Mr-Ap '57. (MLRA 10:5)

1. Chelyabinskiy pedagogicheskiy institut.
(Vegetable gardening—Study and teaching)

APPROVED FOR RELEASE: Thursday, July 27, 2000

CIA-RDP86-00513R00041222

ERN. O.S.

Pupils' experimentation in raising corn. Biol.v shkole no.2:
60-61 Mr-Ap '60. (MIRA 13:8)

1. Chelyabinskiy pedagogicheskiy institut.
(Corn (Maize))

SNESTOPALOVA, T.M.; EIN, O.V.

Ceramic products on a base of clay from new deposit in Transcarpathia. Stroi. mat., det. i izd no. 2115-123 '65
(NTRA 19:1)

1. Lvovskiy filial Gosudarstvennogo nauchno-issledovatel'skogo instituta stroitel'nykh materialov i izdeliy.

R/009/60/000/007/002/003
A124/A026

AUTHOR: Ern, Sergiu, Engineer

TITLE: The Study and the General Diagram for the Selection of Operating
Conditions of Automatic Submerged-Arc Welding 18

PERIODICAL: Metalurgia și Construcția de Mașini, 1960, No. 7, pp. 659 - 663

TEXT: Subject article analyses the main, secondary and accidental elements and phenomena which interfere with the thermo-electric welding process. Some of these elements can be mathematically determined, but the others are difficult to be appreciated. The author also deduces a semi-empiric formula for the value of the welding current, graphically represented in a general diagram, which has to contain all elements necessary for the welding conditions. The main elements of the automatic welding process, i.e., thickness of the sheet in mm (t); area of the welding wire in mm^2 , (q); area of the section of the material deposited in mm^2 , (S); unwinding speed of the wire in m/h (v_d); welding speed in m/h, (v_s); intensity of the welding current in amp, (I); welding tension in v (U); and specific weight of the steel = 7.8 g/cm³ (γ), are shown in Figure 1. Between S, q, v_d and v_s there is a relation, which can be expressed by (1), or for the gen-

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R/009/60/000/007/002/003
A124/A026

The Study and the General Diagram for the Selection of Operating Conditions of Automatic Submerged-Arc Welding

eral diagram by (2). The caloric effect can be computed by the relation

$$K = \frac{U \cdot I}{2.5q \cdot v_d} \quad (7)$$
 Accomplishing weldings of different sections (S) with wires of different thicknesses (q), and measuring U and I , it has been established that the value of K varies between 7.5 and 12, and presents a discontinuity against $q \cdot v_d$ because of the necessity to vary the tension. The K/U ratio presents a continuous variation being a function of $q \cdot v_d$, since I on the other hand is a function of these 2 factors. The welding current (I) is given in case of a d-c current of 140 - 500 amp by the relation (8), and in case of an a-c current of 224 - 800 amp by the relation (9). The d-c current is composed of the interval (10), and the a-c current of the interval (11). The relations (10) and (11) are represented graphically for the selection of the welding conditions. the total quantity of the heat developed is not a continuous proportion with the quantity of the heat necessary for the melting of the wire at 1,400°C. This fact is due to the variation of the welding tension, which is a function of the value of S . The relation $q \cdot v_d = Sv_s$ (12) determines by its left side, accord-

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R/009/60/000/007/002/003
A124/A026

The Study and the General Diagram for the Selection of Operating Conditions of Automatic Submerged-Arc Welding

ing to (10) and (11), the welding current, and by its right side the tension $U = f(S)$. The welding speed (v_s) has no direct influence on current and tension. The control of the welding current is very important. It has been established that a relation exists between t , S and U : $S = f(t)$, and $U = f(S)$. A few suggestions are made for the treatment of the sheets before the welding operation. The quality of the automatically performed welding seams depends on the correct assembly and cleanliness of the sheets. The author then finally presents a diagram for the selection of the welding conditions, which quickly supplies the equilibrium solutions of the seven main elements. The general diagram has been established for the Soviet "TS-17 MU" apparatus, but it can be easily adapted to any other apparatus by notating on the coordinates v_d and v_s the divisions of the respective apparatus. There are 3 tables and 4 figures.

Card 3/3

FERNANDEZ, A.

✓ The influence of prolonged application of fertilizer and of monoculture on the distribution and nitrogen-fixation activity of Azotobacter in sod-podzolized soils. M. V. Pevorov and A. Fernandes. *Izvest. Timiryazev. Sel'skokhoz. Akad.*, 1953, "1957" (Whole No. 8), 145-50.—Prolonged application (more than 40 yrs.) of different combinations of org. and mineral fertilizers show that manure alone or in combination with N-P-K mineral fertilizers stimulates *Azotobacter* activity. Mineral fertilizer alone, especially K salts, depresses the activity. Continuous monoculture also depresses the *Azotobacter* activity. J. S. Josse (1)

ERNA NDES, A.

62. Physiological properties of weakened and atypical Azotobacter cultures grown from nodded podzols. M. V. Fedorov and A. Ernandes. *Mikrobiologiya* 24, 170-9 (1955).— Prolonged culturing of *A. chroococcum* in nodded podzol weakens physiol. activity, especially in N fixation. The enzymes of respiration retain their activity; the N fixation enzymes are inactivated, but are readily reactivated by asparagine, yeast autolyzate, or hay infusion, and also by trace elements such as Mo and B. Lack of these elements may even be the weakening factor in podzols or it may be that podzols contain inhibitors of the growth-stimulating action of yeast autolyzate and trace elements, perhaps due to soil acidity or lack of carbohydrate nutrients. In any case the enzyme system for N fixation is evidently exceptionally sensitive to inactivating influences. J. F. S. (1)

~~GELFAND, I.M.; ERNANDES, L.F.~~

Automatic control of heat-treating furnaces on a programmed
operation. Metallurg 6 no.8:29-31 Ag '61. (MIRA 14:8)

1. Nauchno-issledovatel'skiy institut metiznoy promyshlennosti.
(Furnaces, Heat-treating)
(Automatic control)

ERNAZAROV, E.Yu.

Regeneration of the root system in apple trees. Izv. AN Uz.
SSR no.5:17-24 '56. (MIRA 12:5)
(Regeneration (Botany)) (Roots (Botany)) (Apple)

ERNDT, Aleksander

On synthetic plant growth substance derivatives of p-hydroxy-benzophenone. Roczn. chemii 36 no. 5:921-928 '62.

1. Department of General Chemistry, College of Agriculture,
Krakow.

ACC NR: AP6023858

SOURCE CODE: UR/0108/66/021/007/0044/0051

AUTHOR: Erne Ach (Budapest)

ORG: none

TITLE: Methods of information transmission by the codes that carry information plus address or address only

SOURCE: Radiotekhnika, v. 21, no. 7, 1966, 44-51

TOPIC TAGS: electronic automatic telephone system, information transmission, multichannel telephone system, signal transmission

ABSTRACT: Extreme difficulties involved in synchronizing a large number of channels, in a pulse-code-modulation signal transmission system, are explained. A nonsynchronization system is suggested in which, at the sending end, a code group r_k that has arisen as a result of testing k -th input is supplemented by an "address code"; the latter is a code group p_f that corresponds to the f -th output, the addressee output. The length of the address group depends on the number of channels used; for a 10023-channel trunk, a 19-bit code word is required. Thus, the number of bits is 2.4 times as high as that required for the synchronized system; however, the latter is practically not feasible for such a large number of channels. To reduce the

Card 1/2

UDC: 621.374.372

ERNE, K.

"Sowing maize with soybeans".

p. 59 (Mezhduna Rodnyi Selskokhoziaistvennyi Zhurnal, Vol. 2, No. 2, 1958,
Sofia, Bulgaria).

Monthly Index of East European Accessions (EEAI) LC, Vol. 7, No. 12, Dec. 58.

LIBKOVA, H.; BLASKOVIC, D.; VILCEK, J.; REHACEK, J.; GRESIKOVA, M.;
MACICKA, O., ERNEK, E., MAIER, V.

Incidense of antibodies against tick-borne encephalitis virus in
man and domestic animals in a small village in a natural focus of
infection. J.hyg.epidem., Praha 4 no.3:327-332 '60.

1. Institute of Virology, Czechoslovak Academy of Sciences,
Bratislava.
(ENCEPHALITIS, EPIDEMIC immunol.)

LIBIKOVA, H.; ALBRECHT, P.; ERNEK, E.

Diagnostic horse serum and gamma-globulin against viruses of the tick-borne encephalitis (TE) complex. Acta virol. Engl. Ed. Praha 5 no.4:262 J1 '61.

1. Institute of Virology, Czechoslovak Academy of Sciences, Bratislava.

(ENCEPHALITIS EPIDEMIC immunol) (IMMUNE SERUMS)
(GAMMA GLOBULIN)

NOSEK, J.; REHACEK, J.; ERNEK, E.; GRESIKOVA, M.

The importance of small vertebrates as reservoirs of tick encephalitis viruses in a natural focus in the area of Zlate Moravce. Cesk. epidem. 11 no.6:381-385 N '62.

1. Virologicky ustav CSAV v Bratislave.

(ENCEPHALITIS EPIDEMIC) (ENCEPHALITIS VIRUSES)
(VERTEBRATES)

CZECHOSLOVAKIA

ERNEK, E., MACICKA, O. and ILLES, J. [Virology Institute of CSAV, Bratislava.]

"[Tick-Borne Encephalitis Part] 6. Epizootic Situation Among Domestic Animals."

Bratislava, Biologicke Prace, Vol 8, No 9, 1962; pp 52-58.

Abstract [English summary modified] : Data on contagious diseases in domestic animals in the district Zlate Moravce 1954-1960 and serologic epidemiologic study of Q-fever, brucellosis, toxoplasmosis and leptospirosis. Epizootic conditions are considered favorable to reliable vaccination campaign in this area against tick-borne encephalitis. Three tables.

1/1

CZECHOSLOVAKIA

BLASKOVIC, D., LIBIKOVA, H., ERNEK, E., GRESIKOVA, M., MACICKA, O., VILCEK, J., MAYER, V. and REHACEK, J.; [Virologic Institute of CSAV, Bratislava.]

"[Tick-Borne Encephalitis. Part 9. Planning and Actual Implementation of the Vaccination.]

Bratislava, Biologicke Prace, Vol 8, No 9, 1962; pp 66-75.

Abstract [English summary modified] : Data on serologic diagnosis before and after vaccination in 500 cows, 500 sheep and 500 goats. Both as regards immunogenicity and absolute cost, the live vaccine is superior to the formalized one and the only minor but important advantage of the inactivated one was its safety. Five tables.

1/1

CZECHOSLOVAKIA

GRESIKOVA, M. and ERNEK, E.; [Virology Institute of CSAV, Bratislava.]

[Tick-Borne Encephalitis Part] 11. Transplacental Transmission of the Antibodies in Vaccinated Domestic Animals."

Bratislava, Biologické Práce, Vol 8, No 9, 1962; pp 94-99.

Abstract [English summary modified]: Both live and formalized vaccine against tick-borne encephalitis and louping ill cross the placenta in cattle but the percentage of protected calves varies from 18.6 to 75%. Five tables.

1/1

NOSEK, J.; KOZUCH, O.; LICHARD, M.; ERNEK, E.; ALBRECHT, P.

Experimental infection of the great dormouse (*Glis glis*) with
tick-borne encephalitis virus. Acta virol. 7 no.4:374-376
Jl '63.

1. Institute of Virology, Czechoslovak Academy of Sciences,
Bratislava.
(TICKS) (ENCEPHALITIS)

KOZUCH, O.; NOSEK, J.; ERNEK, E.; LICHARD, M.; ALBRECHT, P.

Persistence of tick-borne encephalitis virus in hibernating hedgehogs and dormice. Acta virol. (Praha)[Eng] 7 no.5:430-433 S '63.

1. Institute of Virology, Czechoslovak Academy of Sciences, Bratislava.

(ENCEPHALITIS, EPIDEMIC) (ZOOSES)
(HIBERNATION)

ERNEK, E.; KOZUCH, O.; LICHARD, M.; NOSEK, J.; ALBRECHT, P.

Experimental infection of *Clethrionomys glareolus* and *Apodemus flavicollis* with tick-borne encephalitis virus. *Acta virol.* (Praha) [Eng] 7 no. 5:434-436 S '63.

1. Institute of Virology, Czechoslovak Academy of Sciences, Bratislava.
(ENCEPHALITIS, EPIDEMIC)

LIBIKOVA, H.; MAYER, V.; REHACEK, J.; KOZUCH, O.; ERNEK, E.;
ALBRECHT, P.; ZEMLA, J.

Study of cytopathic agents isolated from *Ixodes persulcatus*
ticks. *Acta virol. (Praha)* [Eng] 7 no.5:475 S '63.

1. Institute of Virology, Czechoslovak Academy of Sciences,
Bratislava.

(VIRUSES) (TICKS)

CHUMAKOV, M.P.; KARPOVICH, L.G.; SARMAKOVA, Ye.S.; SERGEYEVA, G.I.;
BYCHKOVA, M.V.; TAPUPERE, V.O.; LIEIKOVA, Ye.O.; KAYYER, V.;
RZHEGACHEK, R. [Rehacek, R.]; KOZHUKH, O. [Kozuch, O.]; ERNEK, E.

Isolating from the tick *Ixodes persulcatus* and from sick persons
in Western Siberia a virus differing from the pathogen of tick-
borne encephalitis. Vop. virus. 8 no.1:98-99 Ja-F'63.
(MIRA 16:6)

(VIRUSES) (ENCEPHALITIS—MICROBIOLOGY)

LIBIKOVA, H.; GRESIKOVA, M.; REHACEK, J.; ERNEK, E.; NOSEK, J.

Immunological surveys on natural foci of tick encephalitis.
Bratisl. lek. listy 43 no.1:40-53 '63.

1. Virologicky ustav CSAV v Bratislave, riaditeľ akademik
D. Blaskovic.

(ENCEPHALITIS, EPIDEMIC)
(ARBORVIRUS INFECTIONS)
(NEUTRALIZATION TESTS)
(ANTIBODIES)

LIBIKOVÁ, H., REHÁČEK, J., MAÝER, J., KOZULÍK, O., EKHÉŘE, *_____*

Tele-borne external ticks removed by multiple methods
from 1000 patients and 1000 dogs, especially from cattle
and sheep.

To: Institute of Veterinary, Faculty of Veterinary Medicine, Štefánikova,
Bratislava.

†

LIBIKOVA, H., REHACEK, J.; GRESIKOVA, M.; KOZUCH, O.; SOMOGYIOVA, J.
Ernek, E.

Cytopathic viruses isolated from ixodes ricinus ticks in
Czechoslovakia. Acta virol (Praha) [Engl] 8 no.1:96 Ja'64.

1. Institute of Virology, Czechoslovak Academy of Sciences,
Bratislava.

*

LIBIKOVA, H.; MAYER, V.; KOZUCH, O.; REHACEK, J.; FRNEK, E.; ALBRECHT, P.

Isolation from *Ixodes persulcatus* ticks of cytopathic agents
(Kemerovo virus) differing from tick-borne encephalitis virus
and some of their properties. *Acta virol. (Praha)* [Eng.] 8
no.4:289-301 Jl '64.

1. Institute of Virology, Czechoslovak Academy of Sciences,
Bratislava.

ERNEK, E.; LICHARD, M.

Role of the English sparrow (*Passer domesticus*) in the circulation of tick-borne encephalitis virus. *J. hyg. epidem. (Praha)* 8 no. 3:375-379 '64

1. Institute of Virology, Czechoslovak Academy of Sciences, Bratislava.

GRASZ, J.A., M.; KUCERA, J.; KLAUCH, J.; KUDLA, V.; LICHNER, M.

Study on the ecology of Tribes virus. Acta virol. (Praga)
[Eng.] 9 no.1863-28 Ja 1965

1. Institute of Virology, Czechoslovak Academy of Sciences,
Bratislava.

ERNEY, Gyorgy

New nomenclature of gear wheels, Szabvany kozl 13 no.7:158-160
Jl '61.

ERNEY, Gyorgy, okleveles gépész mérnök

Comparing the precision standards of up-to-date cylindrical cogwheels. Gap 16 no. 118415-424 N '64.

LEND'YEL, V.I.; ERNEST, B.M.

Use of analyticity conditions of the scattering amplitude
in determining the coupling constant. Dokl. i soob. UzhGU.
Ser. fiz.-mat. i ist. nauk no.5:14-16 '62. (MIRA 17:9)

JELEA, Al.; ERNEST, Ilie; PIRVU, V.; NUTA, M.; DIACONU, J.

Contributions to the study of trypsin treatment in bronchopulmonary
disease. Rumanian med. rev. no.2:25-28 '62.
(TRYPSIN) (LUNG DISEASES)

JELEA, Al., dr.; ERNEST, Ilio, dr.; PIRVU, V., dr.; NUTA, M., dr.; DIAGONU, J.,
intern

Contributions to the study of trypsin therapy in bronchopulmonary
diseases. Med. intern. 14 no.1:67-72 Ja '62.

1. Lucrare efectuata in Institutul de medicina interna al Academiei
R.P.R. si M.S.P.S., director: acad. N.Gh. Lupu.
(LUNG DISEASES therapy) (BRONCHI diseases)
(TRYPSINS therapy)

CAPEK, A.; TADRA, M.; KAKAC, B.; ERNEST, I.; FROTIVA, M.

Microbiological transformation of derivatives of hexahydronaphthalene acid. Folia microbiol. 7 no.4:253-254 '62.

1. Institute of Pharmacy and Biochemistry, Prague 3.
(NAPHTHALENES - metabolism) (LACTONES - metabolism)
(FUNGI - metabolism) (ACTINOMYCES - metabolism)

ERNEST, I; JÍLEK, O; VEJDĚLEK, Z; PROTIVA, M.
Czechoslovakia

Research Institute of Pharmacy and Biochemistry -- Prague
- (for all)

Prague, Collection of Czechoslovak Chemical Communications,
No 4, 1963, pp 1022-1029

"Synthetic Experiments in the Group of Hypotensively
Active Alkloides XXVI. On Some New (-)-Methyl-
Reserpate-Ester."

ERNEST, I.; KAKAC, B.; PROTIVA, M.

Synthetic experiments in the group of active hypotensive alkaloids. Pt.31. Coll Cz Chem 29 no.1:251-265 Ja'64.

1. Forschungsinstitut fur Pharmazie und Biochemie, Prag.

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3-Substituted quinuclidines. III R. Lukes and J. Ernest Chevre Polytech., Prague, Collection Czech. J. Chem., 1972, 15, 150-5 (Brockmann, et al., 1944, 5,213). 3-Acetoxyethyl-3-(bromomethyl)quinuclidine. Melts (D), after hydrolysis with 0.5% HBr in MeOH, distn of the MeOH, boiling with aq. AgO, and evapn to dryness with AgO, gave 1-methyl-3-methylenquinuclidinium picrate (II), decomps 257-8°, and a small amt. of 3-carbethoxy-3-methoxy-1-methylquinuclidinium picrate (III), m. 213-14°. III was not isolated until after II was converted to the chloride (addn of HCl, filtration, and evapn with Pt(NO₃)₄ and hydrogenated PtO₂, 910 mm, 25°, in EtOH) to 1-(bromomethyl)quinuclidinium picrate (IV), decomps 200°. IV acetate (prepd. from the picrate via the chloride and AgOAc) decomps. at 100-200° to 3-methylquinuclidine (V), b. 170-82°; picrate, m. 225°; chloroplatinate, m. 218.5. I formed a different sample (VI) of II, decomps. 261°, on boiling with Ba(OH)₂, pptn. of the Ba with CO₂ and H₂S₂O₃, shaking with AgO, acidification with HOAc, and vacuum evapn. The acetate related to II (prepd. via the chloride and hydrolyde) formed on pyrolysis 3-methylenquinuclidine (VII), bns 170-1°, μ 1.4006, δ 0.0100, picrate, m. 200°; chloroplatinate, decomps. 101°. Hydrogenation of VII (PtO₂, 910 mm, 25°, in HOAc) formed V.

J. H. Scott

cA ERNEST, I.

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Proof of the preparation of 3-vinylquinuclidine. *Ernest (Collection Czechoslov. Chem. Commun. 15, 329-34 (1950) French).* The synthesis of 3-(2-hydroxyethyl)quinuclidine (I) from 3-ethyl-3-hydroxyquinuclidine (II) is described. Dehydration of I produces 3-ethylquinuclidine rather than the 3-hydroxy deriv. To NaClO_4 (from 26 g. Na and CaH_2 in 300 cc. liquid NH_3) was added 29 g. quinuclidine in 80 cc. Et_2O , and after evapn. of the NH_3 and decomprn. with 200 cc. acidified ice water, the product was made alk. and extd. with CHCl_3 , giving on removal of the solvent, 20.2 g. colorless crystals of II, m. 101.2°; *picrate*, m. 170.0.5°. II (180 mg.) in 10 cc. EtOH hydrogenated in the presence of 50 mg. 3% Pd-CaCO_3 at atm. pressure, yielded 3-ethyl-3-hydroxyquinuclidine (III), m. 110.5°; *picrate*, m. 150°. Reduction of 28.5 g. II in 500 cc. EtOH with 1.30 cc. H_2 at atm. pressure over 2.5 g. 3% Pd-CaCO_3 , removal of the solvent, and sublimation of the crystall. residue gave 27.5 g. 3-ethyl-3-hydroxyquinuclidine (IV), m. 90.0.5°; *picrate*, m. 151.8°. IV (29.0 g.) was heated in a sealed tube with a 10% soln. of HBr in AcOH for 2 hrs. at 60.5°, the solvent removed *vacuo*, the residual syrup heated 2 hrs. with 100 g. KOH in 750 cc. AcOH , the KBr and AcOH removed, the residue refluxed with 1500 cc. 10% KOH-MeOH , the solvent removed, the residue dissolved in H_2O , extd. with CHCl_3 , the ext. dried, evapd., and the residual syrup (27.8 g.) dissolved in CaH_2 and chromatographed over Al_2O_3 , giving α -[3-(2-hydroxyethyl)dihydroquinuclidine] (V), m. 91.5.5° (*picrate*, m. 121.3.5°), and β -isomer (VI), m. 18.8.5° (*picrate*, m. 151.2°). V (8.4 g.) in 100 cc. EtOH absorbed 703 cc. H_2 in 2 hrs. in the presence of 0.2 g. Pd-CaCO_3 and gave 5.1 g. I, m. 121.0°, m. 31.0°; *picrate*, m. 91.5.5°. VI (7.8 g.) in 100 cc. EtOH and 0.2 g. Pd-CaCO_3 absorbed 1160 cc. H_2 and gave 7.4 g. I, m. 118.0°.

(1.05 g.) in 10 cc. PhMe was refluxed 30 min. with 10 g. Pd-CaCO_3 in PhMe added dropwise, the solvent removed, the residue heated 2 hrs. at 100.0°, decomprn. with H_2O , the remainder of the solvent steam distd., the residue made alk. and the base steam distd., from the HCl salt of the diolate the *picrate*, m. 128.5.0°, of 3-ethyl-3-hydroxyquinuclidine (VII) was prep'd.; *chloroplatinate*, m. 150.80°, I (0.36 g.) heated 3 hrs. at 180° with 10 cc. concd. HCl in a sealed tube also gave VI, I (0.35 g.) heated 1 hrs. at 115° in a sealed tube with 10 cc. concd. HCl gave 3-(2-hydroxyethylquinuclidine (VIII) isolated as the *picrate*, m. 130.5.7°. I (150 mg.) in 5cc. xylene was shaken with 135 mg. K dust in 3 cc. xylene, heated 3 hrs. on an oil bath, the absolute suspension shaken 30 min. with 100 cc. freshly distd. $\text{C}_2\text{H}_5\text{OH}$, and 200 mg. MeI added, the mixt. stirred several hrs., then heated 6 hrs., and the brown ppt. collected, washed with Et_2O , and decomprd. in a Hückel oil still, giving 200 mg. of distillate, which after purification was converted to the *picrate*, m. 101.5.2°, of the *N*-methylchloroplatinate of 3-ethylquinuclidine. $\text{CaH}_2\text{-NaS}_2$ (1.00 g.) in 100 cc. EtOH solution in AcOH was heated 15 hrs. at 100° in a pressure bottle to give 7.7° *bromomethylquinuclidine* (II), m. 107.8°; *picrate*, m. 118.0°. VII (1.02 g.) refluxed 5 hrs. with 170 cc. 10% KOH in MeOH , steam distd., and acidified with HCl and H_2O acid gave the 3-(2-methoxyethylquinuclidine *picrate*, m. 128.0°.

Bernard Klein

1951

C. R. ERNEST, I.
1951

Organic Chemistry
10

Synthesis of 3-vinylquinuclidine. I. Ernest (Tech. Univ., Prague). *Collection Czech. Chem. Commun.* 19, 490-93 (1950) (in English); cf. *C.A.* 45, 3818c. — When 3-(2-hydroxyethyl)quinuclidine (*C.A.* 45, 3848c) is heated 30 min. with α -C₆H₅(CO)NO and PhSO₂H, 3-ethylidenequinuclidine (I) alone is formed, but if the heating is shorter, a mixt. of I and 3-vinylquinuclidine (II) results. These were sep'd. by fractional cryst. of the styphnates. The picrates formed an isomorphous mixt. II, liberated from the acid styphnate, bp 98° (bath temp.), d₄²⁰ 0.9005, n_D²⁰ 1.6990; picrate, m. 150° 1²; acid styphnate, C₂₁H₁₆N₄O₄, m. 152°; *infrared* spectra given.

Alfred Hollman

ERNEST, I.
CA

16

Decomposition of bis(diazo ketones) with cupric oxide, Ivan, Károly and Jiri Hofman (Tech. Univ., Prague, Czech.), *Chem. Listy* 65, 201-4 (1981).—[Bis(diazo ketones) give, on decomp. with CuO, unsat'd cyclic dithiones. The dichloride (I) of 2-carboxycyclopentanone-1,1-dicarbonyl (IA) was precip. by refluxing 7.2 g. IA 4 hrs. with 20 g. CuCl in 25 ml. xylene, stripping off the xylene and PCl_3 in vacuo, and distg. the crude I (6.7 g. b.p. 77-112°, 1.6 g. b.p. 114-37°); residue yielded 3.8 g. I, b.p. 70°. With Bz_2Cl , IA gave mostly the anhydride, b.p. 114-15°, which yielded I with PCl_3 , $(\text{CH}_3\text{COCl})_2$, $\text{CH}_3(\text{CH}_2\text{COCl})_2$, and I treated with a 6-mol. excess of CH_3I_2 in ether gave the bis(diazo ketones), 1,1-bis(2,2-butanedione) (II), 1,7-bis(iso-2,2,2-heptanedione (III), and 1-diazoisopropyl-2-(2-diazoacetoxy)cyclopentane, probably the *cis*-isomer (IV), resp. II could not be isolated, by distg. off the ether (it decomposed even at room temp.), but its m.p. in CaH_2 was stable. III (0.1 g.) from 10 g. $\text{CH}_3(\text{CH}_2\text{COCl})_2$, yellow needles from ether, m.p. 63-45°, decompr. at 108°. IV, yellow oil, decompr.

above 100°, stable in *CCl₄*. II (3 g.) with HCl in 300 ml. ether gave 1,6-dichloro-2,6-benzoquinone (V), white crystals, m. 87.5-8° (from *EtOH*). III (2 g.) gave similarly 2 g. crystals in 73.5° (70° after recryst., from 1:2 *EtOH*-ether mixt.). III (1 g.) treated with 13 ml. 30% NaOH in 40 ml. dioxane at 60-70° in the presence of 2.3 ml. 10% *AgNO₃*, and the dioxane dried off after 45 min. gave 0.4 g. pinacolambone, m. 170-173° (174° A° after recryst., from *EtOH*), hydrolyzed to the acid. IV (0.55 g.) with HCl in 70 ml. ether yielded 0.45 g. 2,6-dichloro-1-phenylcyclo-

(3-chloroacetyl)cyclopentanone (VII), colorless liquid, bp. 124-5°; *bd*(3,4-dinitrophenylhydrazine), reddish crystals, m. 176-7°, yield 2,4-dinitrophenylhydrazone. The hydrazine ketones I were decomposed, by refluxing with equal amounts of CuO in Cells, III (prep'd. from 9 g. (CH₃COCl)₂) in 2 l. Cells was refluxed 3 hrs. with 9 g. CuO, the solution filtered, evaporated, to 50 ml., the resinous material removed, and the filtrate chromatographed over Al₂O₃, yielding 0.36 g. crystal V (Cells fraction) and 3 undecarboxylated fractions in ether (0.073 g.) and MeOH (0.72) g.). No cyclic ketone was obtained. III (4 g.) similarly gave 0.116 g. Cells fraction and 0.170 g. yellow ether fraction. The Cells fraction yielded the bis(RuOH-CH₂N)₂ of 2-cyclopentene-1,4-dione (decarboxyl.) (from compn. of (CH₃Cl)₂CO₂NH₂) prep'd. from (CH₃Cl)₂CO₂NH₂ (COCl) gave no individual ketone on chromatography. IV (prep'd. from 2.8 g. I) in 250 ml. Cells was refluxed 3 hrs. with 3 g. CuO, the CuO removed, the solution evaporated to 20 ml., treated with 50 ml. ether, the resinous ppt. filtered off, the filtrate evaporated, and the residue extracted with ether, yielding 1 g. of a syrupy material which on chromatography over Al₂O₃ No. 2 gave 0.1 g. Cells fraction and 0.060 g. ether fraction (no individual ketone). The Cells fraction yielded by another chromatography over Al₂O₃ 0.167 g. of bis(cyclopent-3,4-diene-2,3-dione). *bd*(3,4-dinitrophenylhydrazine), m. 231° (decarboxyl.) (from RuO₄).

CA

ERNEST, I.

Allylic rearrangement. Ivan Ernest, (Tech. Univ.,
Prague, Czech.). *Chem. Listy* 46, 58-61 (1952) --A review
with 73 references. M. Hudlicky

Preparation of *p*-nitroacetophenone. J. Erényi and Z. Verely (Vysoká škola chem., Prague, CZECHOSLOVAKIA 1977, 740-81083). *p*-O₂NCH₂CH₂Cl (I) has been prep'd. from PhCH₂XMe (II) (where X = Cl or Br) in an over-all yield of 28% by the following sequence of reactions: II obtained by the addn. of HCl or HBr to PhCH₂CH₃, gave, by nitration, *p*-O₂NCH₂CH(ONO₂)Me (III), which was transformed to *p*-O₂NCH₂CH(OAc)Me (IV). IV hydrolyzed to *p*-O₂NCH₂CHMeOH (V), and oxidized to I. Direct conversion of III to I was not successful, and oxidation of IV to I gave low yields. *p*-O₂NCH₂CH(ONO₂)CH₂Br (VI) was prep'd. like III by the ulturation of PhCH₂Br. m. 48-0° (the yields for X = Cl and Br were 39.1 and 42% resp.). Refluxing 1 g. III, 2 g. NaOAc, 2 g. urea, and 2 g. AcOH in 30 ml. Ac₂O 4 hrs., distg. off the volatile compds. *in vacuo*, treating the residue with H₂O, extg. with Et₂O, and evapn. the exts. gave 0.8 g. (50%) IV, m. 65-6°. V (*p*-nitroacetate, m. 137-7.8°), obtained from IV, oxidized at 20° with dil. chromic mxt., yielded 87% I, m. 78.5-8°. IV (22 g.) heated 2 hrs. at 60° with 10 g. K₂Cr₂O₇, 21.5 ml. concd. H₂SO₄ and 68 ml. H₂O gave, 50.5% I. VII (30 g.) added to 300 g. HNO₃ (d. 1.8) at -5° gave 17 g. (63%) VII, m. 73.5-8.7° (from Et₂O). M. Hadlicky

✓ Herout, V., Keil, B., Protiva, M., Hudlicky, M., Ernest,
and Gut, J.: Laboratorní technika organické chemie
Prague: Nakl. CSAV, 1954. 756 pp. Kčs 80. Re-
viewed in Chem. Listy 49, 1415(1955).

Chem) Educ

Herout, V., Keil, B., Protiva, M., Hudlicky, M., Ernest,
Organic Chemistry Laboratory Techniques. Publishing House CSAV. 1954.
756pp. Kčs. 80. Reviewed in Chem. Listy 49, 1415(1955).

PM

ERNEST, IVAN

CZECH

Decomposition of diaz ketones by cupric oxide. II
 Decomposition of monoacetylcarboxylic esters. A new
 method for the preparation of higher paraffin- α,ω -dicar-
 boxylic acids. Ivan Ernest (Vysoká škola chem.-
 techn. Praha). *Chem. Listy* 48, 847-57 (1954); *Collection
 Czechoslov. Chem. Commun.* 19, 1176-80 (1954) (in
 German); *J. C. S. 46, 7048; -CICO(CH₂)_nCO₂Et (I)*
 $n = 0, 1, 2, 3, 4$, were transformed to N(CHCO(CH₂)_n)
 CO₂Et (II) the decompn. of which with CuO gave -CHCO₂
 (CH₂)_nCO₂Et (III). Hydrogenation of III ($n = 2, 3, 4, 5$)
 yielded [CH₂CO(CH₂)_nCO₂Et] (IV), which were hydrolyzed
 to [CH₂CO(CH₂)_nCO₂H] (V). Three of the IV ($n =$
 $1, 2, 3$) were transfo. to the corresponding [CH₂C(SCH₂)
 CH₂CO(CH₂)_nCO₂Et] (VI) the desulfurization of which with
 Raney Ni gave [CH₂CH₂(CH₂)_nCO₂Et] (VII). [CH₂CH₂
 (CH₂)_nCO₂H]₂ (VIII) were obtained from VII by alk. hy-
 drolysis. Refluxing V ($n = 3$) 60 min. with 25% wt. NH₃
 gave pyrrol- α,α' -divaleric acid (IX), m. 153-00.5° (from
 H₂O). I were prep'd. by heating the corresponding add-
 esters with excess SOCl₂ (n , and b.p. are given): 0, 129.6-
 30.5°; 1, b₁ 60-60.5°; 2, b₂ 80-80°; 3, b₃ 107.5-3.5°; 4,
 b₄ 117°; 5, b₅ 117-12°; 6, b₆ 148-51°. Adding 0.1 mole I in
 200 ml. Et₂O during 60 min. at 0° to a soln. of 0.3-0.36
 mole CH₂N₂ (prep'd. from 42 g. NH₂CONMeNO) in 500
 ml. Et₂O, allowing to stand 60-90 min. at room temp., and
 distg. off Et₂O *in vacuo*, at 30-40° yielded Et₂O soln. of II.
 N(CHCO₂)₂ was isolated in 69% yield by cooling the
 soln. (evap'd. to 200 ml. vol.) to -15°, m. 72.5-4°. The
 homologous II were evap'd., dissolved in Et₂O to 50-70 ml.,
 the solns.稀d. with 100-150 ml. C₆H₆, and distg. *in vacuo*
 to the original vol. Repeating this action substituted C₆H₆
 for Et₂O, and the solns. thus obtained were used for the

reactions. Pure II obtained by distg. off the Et₂O₂ are unstable, orange liquids. Refluxing 0.1 mole II in 250 ml. C₆H₆ with 5 g. powdered CuO 20-30 min. (or 0.2 g., 60-60 min.), filtering off the CuO, and evapg. the solvent *in vacuo* gave crude III mostly cryst. (except for n = 3). First crops were obtained by filtration, addnl. crops by chromatography of the mother liquors. Results of the prepns. of III are as follows: n, yield (based on II) in %, m.p. (from petr. ether): 0, 2.2, 133-4.5%; 1, 0.9, 100-7° (from C₆H₆-Et₂O(1:1); 2, 20.8, 62-3° (deirime, m. 111-12°), 3, 62.0, 35-5.5° (bis-*p*-nitrophenylhydrazene, m. 221, decompn.); 4, 45.4, 51-8.3° (deirime, m. 118°; bis-*p*-nitrophenylhydrazene, m. 161° (decompn.); 5, 23.1, 50.5-2°; 6, 32, 74-5.5° (bis-*p*-nitrophenylhydrazene, m. 140-2°, decompn.). III was hydrogenated in EtOH-C₆H₆ soln. at 1 atm., 20°, and 5% Pd-CaCO₃. 1.14-1.24 moles H per mole II was consumed. The following IV are prepnd. (n, yields in %, and m.p. given): 3, 71.8, 20°; 4, 65.8, 50.5-3°; 5, 60.7, 40.5-8°; 6, 84.6, 67-9°. Heating IV with concn. HCl 60-90 min. at 100° (the ester n = 8 was refluxed 2 hrs. with HCl) gave V (m.p.): 2, 155-0°, 3, 134-5°, 4, 123-9.5°, 5, 132.3-3.5°, 6, 133-4.5°. Treatment of IV with (CH₃Si)₂N₂, Na₂SO₄, and ZnCl₂ in a dioxane soln. yielded VI (n, m.p.): 4, 60-1.5° (64.0%); 5, 30.5-0°; 6, 9, liquid. Refluxing VI with ten fold amt. of Raney Ni 8 hrs. yielded VII (n, yield in %, m.p.): 4, 84.2 (based on VI), 27.5-9°; 5, 53.5 (based on VI), 37-9°; 6, 72.0 (based on V), 61-2.5°. Free acids VIII obtained by sapon. with 10% aq.-alc. NaOH were crystd. from C₆H₆ (n, m.p.): 4, 125-0°, 5, 124.5-5°, 6, 125.5-0.5°. M. Hudlicky

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HNEVSOVA, V., SNELY, V., ERNST, J.

Hydrogenated at 50° with PbO in C_6H_6 yielded 2.03 g. of *di-Me₂isobutylidene-2,2-dione-1,3-diolate* (X), m.p. 107.5-108° (from $MeOH-C_6H_6$), IX (1.52 g.) was converted by treatment with 2.6 ml. (18.61 g.) and 1.5 ml. $Hg₂Et₂O$ to the corresponding *bis[carboxymercaptoether]* (XI), m.p. 77°. Desulfurization by boiling in C_6H_6-MeOH (1:1) gave 0.77 g. of a product m.p. 93-94° which on treatment with Raney Ni (V) gave 883 mg. of a product m.p. 76-77° (from $Et_2O-C_6H_6$), m.p. 100-102° (from $CHCl_3-MeOH$). X was also obtained in 70% yield by a shorter alternative synthesis including treatment of 600 mg. VIII with $(HSCH_2)_2$ and $Bf_3\cdot Et_2O$, subsequent desulfurization and hydrogenation. Alk. hydrolysis of *di-Me₂isobutylidene-2,2-diolate* (X), m.p. 128.5-91.5°, similarly, treatment of *Me₂chloro-2,2-dihydroxy-3,3-dimethyl-1,1-dioxane-4-carboxylate* (XII) (77.8% yield, from the *Me₂ester chloride* of succinic acid and $C_6H_5N_3$) with CuO gave *di-Me₂isobutylidene-2,2-diolate* (XI), m.p. 120-121.5°, in 33.5% yield, while *Me₂chloro-2,2-dihydroxy-3,3-dimethyl-1,1-dioxane-4-carboxylate*, *Me₂chloro-2,2-dihydroxy-3,3-dimethyl-1,1-dioxane-4-carboxylic acid*, and CH_3N_3 was converted to *di-Me₂isobutylidene-2,2-dione-1,1-diolate*, m.p. 122-123°, in 22.2% yield. IV. Substitution of unsaturated *di-Me₂isobutylidene-2,2-diolate* (X) by *Me₂chloro-2,2-dihydroxy-3,3-dimethyl-1,1-dioxane-4-carboxylate* (XII) (100% conversion) on the activity of the used $PbCl_2\cdot 2H_2O$ gave 1.4 g. of a product m.p. 120-121°. After 3 hr. the mixture was shaken with C_6H_6 and the *Me₂chloro-2,2-dihydroxy-3,3-dimethyl-1,1-dioxane-4-carboxylate* (XII) was removed by vacuum distillation, m.p. 121-122°, in 1.6 g. (65% yield) of *Me₂isobutylidene-2,2-diolate* (X). No Me_2CO or CH_3CO_2Et gave 1.4 g. of *Me₂isobutylidene-2,2-diolate* (X). No Me_2CO or CH_3CO_2Et gave 130 mg. of *Me₂isobutylidene-2,2-diolate* (X), m.p. 135-136°, in 27% yield. Alk. hydrolysis of XI gave the free acid (XI), m.p. 125-126°, in 1.6 g. (65% yield).

(continued)

HANESOVÁ, V., SMELY, V., ERNST, I.

that has been deactivated by boiling 2 hrs. with MgCO_3 gave a fraction which was identified as a mixt. of 59% II and 41% of the corresponding unsatd. ester, probably *di-Et-dec-6-ene-1,12-dicarboxylate* (IV), characterized by coulometric analysis and by hydrogenation, yielding II. When a 12-hr. inactivation was used, deactivation of 3.5 g. I gave a 669-mg. fraction, $\text{b}_{10} 135-45^\circ$, which on alk. hydrolysis yielded crystals, m. 125-7°, probably of $\text{HOOC(CH}_2\text{CH}_2\text{CH}_2\text{CHCO(CH}_2\text{)_2\text{CO}_2\text{H}}$, whereas a 6-hr. inactivated catalyst produced a fraction, $\text{b}_{10} 133-7^\circ$, apparently of IV, identified by hydrogenation which gave II and after alk. hydrolysis yielded III. A parallel expt. from 4.7 g. I gave a 1.18-g. fraction, $\text{b}_{10} 124-7^\circ$, which was chromatographed on Al_2O_3 yielding by alk. hydrolysis of the ligroine eluate 40 mg. cryst. *deca-6-ene-1,12-dicarboxylic acid*, m. 107-9°, confirmed by coulometric analysis. Attempts were made at overcoming difficulties encountered in the prepn. of unsatd. dicarboxylic acids of the type $\text{RO}_2\text{C}(\text{CF}_3)_2\text{COCH}_2\text{CH}_2\text{CO}_2\text{R}$ (V) by prep. addn. compd. of V with anthracene (VI), however, without success. The adduct of Ia and VI obtained by heating 5 hrs. powd. mixt. of 3.0 g. VI with 6.8 g. V ($n = 6$, R = Et) forms crystals, m. 78-9° (from cyclohexane-C₆H₆), yielding on sapon. crystals, m. 183-4° (from C₆H₆-AcOH). Similarly was prep'd. the adduct of *di-Me-cet-4-eno-3,6-dione-1,6-dicarboxylate* with VI from 0.7 g. VI and 1.0 g. V ($n = 2$, R = Me), forming needles, m. 130-5° (from C₆H₆), and yielding on sapon. crystals m. 213-14° (decompn.) (from AcOH).

L. J. Urbanek

3/2

CZECHOSLOVAKIA / Organic Chemistry. Synthetic Organic Chemistry

G-2

Abs Jour : Ref. Zhur. Khimiya, No 3, 1950, 7975

Abstract : 29.2 gm ethyl oxalate and KOC_2H_5 (out of 7.02 gm of K and 8.65 gm of alcohol in 85 ml of ether). The benzylisothio salt of II has a m.p. of $148^{\circ}C$ (in CH_3OH -ether). II and an alcoholic solution of HCl (~ 20 C, 2 days) react to yield ethyl β -(2-pyridyl)- α , β -dichlorovalerate, b.p. $136^{\circ}C/1$ mm Hg, $n^{20}D$ 1.4842. By hydrogenating the Ba salt of II in an aqueous solution at $25^{\circ}C$ and 730 mm Hg over PtO_2 , β -(2-pyridyl)- α -oxyvaleric acid (III), m.p. $137-139^{\circ}C$ (in alcohol) was prepared; the ethyl ester of the latter acid had a b.p. of $150-155^{\circ}C$ (bath temperature)/1 mm Hg, $n^{20}D$ 1.5040. A mixture of both stereoisomers of ethyl β -(2-pyridyl)- α -oxyvalerate, b.p. $123-129^{\circ}C/2.5$ mm Hg, m.p. $66-80^{\circ}C$ (in petroleum ether) was separated by hydrogenating free II (water, PtO_2 , $19^{\circ}C$, 965 mm Hg) and esterifying the product with alcoholic HCl. The racemate (m.p. $92-93^{\circ}C$) was

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ERNEST, I.; PITHA, J.

SCIENCE

Periodical COLLECTION OF CZECHOSLOVAK CHEMICAL COMMUNICATIONS. SBORNÍK CHEMKOSLOVATSKÝKH KHIMICHESKÝKH RABOT. Vol. 23, no. 1, Jan. 1958.

ERNEST, I.; PITHA, J. Quinolizidine dérivatives. I. Catalytic hydrogenation of δ -(2-pridyl)- α -oxovaleric acid. In German. p. 125.

Monthly List of East European Accessions (EEAI) LC, Vol. 8, no. 3, March, 1959. Uncl.

CZECHOSLOVAKIA/Organic Chemistry. Synthetic Organic Chemistry. G

Abs Jour: Ref Zhur-Khim., No 2, 1959, 4619.

CuO, are cyclized by the action of strong acids in CH₃COOH medium to form unsaturated 2,5-disubstituted derivatives of furan of the type $\text{OCR}=\text{CHCH}=\text{CCH}=\text{CHR}'$. The reaction in all probability proceeds by a mechanism similar to that of the opening of the furan ring according to Marckwald. Preparation: 60 gms of butyl chloride on treatment with diazo-methane in ether solution at -20° give 1-diazo-pentanone-2; the ether is distilled off and the product is decomposed by refluxing for 15 min with 6 gms CuO in 2 liters of C₆H₆, giving 5-decene-4,7-dione (I), yield 28.5%, mp 55-56.5° (from CH₃OH). Using a similar procedure, dihydrocinnaryl chloride gives a 27.3% yield of 1,8-diphenyl-4-octene-3,6-dione (II), mp 85-85.5° (from alc). 2 gms of the methyl

Card : 2/9

CZECHOSLOVAKIA/Organic Chemistry. Synthetic Organic Chemistry. G

Abs Jour: Ref Zhur-Khim., No 2, 1959, 4619.

chromatography on Al_2O_3 (elution with benzene); the free acid (Mp 101°; from aqueous alcohol) on oxidation with $KMnO_4$ gives suberic acid and sebamic acid. I yields 2-propenyl-5-propylfuran, bp 47-50°/0.4 mm, n^{20}_D 1.5008; the methyl ester of 4-heptene-3,6-dione-1-carboxylic acid gives 5-methyl-2-(β -carboxethoxyvinyl)-furan (IV), yield 74%, mp 36-37°, bp 65-70°/2mm; the free acid (V) has an mp of 154° (from water). 5-ethyl-2-(β -carboxethoxyvinyl)-furan (VI), mp 47-48°, bp 75-80°/1.5mm, was also synthesized from the methyl ester of 4-octene-3,6-dione-1-carboxylic acid. The UV spectra of III-VI are given.

VI. Esters of asymmetric unsaturated diketocarboxylic acids for the syntheses reported in the preceding

Card : 4/9

CZECHOSLOVAKIA/Organic Chemistry. Synthetic Organic Chemistry. G

Abs Jour: Ref Zhur-Khim., No 2, 1959, 4619.

fold excess of VIII is used, the yield [sic] is increased to 33%. Using a similar procedure, VII and 1-diazo-2-butanone give the methyl ester of 4-octene-3,6-dione-1-carboxylic acid, yield 15.5%, bp 125-130°/2mm, mp 47-48° (from petroleum ether) (the product was separated by distillation after the removal of 1,2-dipropionyl ethylene, yield 53%, bp 80-85°/3mm, mp 52-53° (from petroleum ether)); VIII and the methyl ester of ω -diazoacetylvalerianic acid after distillation of IX (47%) and crystallization of the methyl ester of dodecene-6,5,8-dione-1,12-dicarboxylic acid (yield 14%, mp 93-94° (from CH_3OH)) give the methyl ester of 6-nonene-5,8-dione-1-carboxylic acid, yield 21%, bp 120-140°/2 mm, mp 53-54° (from petroleum ether).

Card : 6/9

10

CZECHOSLOVAKIA/Organic Chemistry. Synthetic Organic Chemistry. G

Abs Jour: Ref Zhur-Khim., No 2, 1959, 4619.

benzoic acid (X), yield 98%, mp 70-71°; the ethyl ester of p-dinzoacetylbenzoic acid (XI) yield 99%, mp 65-66°. The following compounds were prepared by refluxing X or XI for 15 min with CuO in C₆H₆: 33.5% 1,2-bis-(*n*-carbethoxybenzoyl)-ethylene, mp 131-132° (from ethyl acetate) (XII); 36.5% 1,2-bis-(*p*-carbethoxybenzoyl)-ethylene (XIII), mp 194-195° (from benzene). 1,2-bis-(*o*-carbethoxy-benzoyl)-ethylene, mp 160° (decomp; from alc-benzene) was obtained by a similar procedure from the acid ethyl ester of phthalic acid without the separation of intermediate products. The hydrogenation of XIII to XII over Pt (from PtO₂) at 22° and at normal pressure yields 1,2-bis-(*p*-carbethoxybenzoyl)-ethane, mp 157-158° (from alc); similarly 1,4-bis-

Card : 8/9

11

SEARCHED	1
INDEXED	1
ABSTRACTED	1
FILED	1
AUTHOR	J. Jithé, Y. Grunek
INIST.	1
TITLE	Synthetic Approaches to the Alkaloids. Part VI. Isolation of the crystalline γ -(2-oxopropyl)- β -conopeptides.
ORIG. PUBL.	Chem. Lett., 1979, p. No. 16, 3627-3630
ABSTRACT	The hydroxylation of γ -(2-oxopropyl)- β -conopeptides, non-2 (I) and 2-(2-oxopropyl)- β -conopeptides, non-2 (II), is found to be selectively at the 3-position. The structure of the 3-hydroxy- β -conopeptides is determined by the position of the C-terminal carboxyl group (II'). On the basis of mutual correlation, in view of the chromatographic data, the authors assume that the hydroxy group is at the C(10) position. The structure of the 3-hydroxy- β -conopeptides is determined by the position of the C-terminal carboxyl group (II'). On the basis of mutual correlation, in view of the chromatographic data, the authors assume that the hydroxy group is at the C(10) position.

CARD:

1/7

C-37

1. JOURNAL	:
2. LITERATURE	:
3. SOC. JOUR.	: RZKohl., No. 23 1959, No. 407
4. AUTHOR	:
5. TITLE	:
6. PUBL.	:
7. CITE. PUB.	:
8. ABSTRACT	: In this paper, from the products of hydrolysis of 5 -(pyridyl-2)- α -ketovaleric acid (V), it was possible to prepare a small quantity of 2-chinolysimidocarbonylic acid (VI), identical with the acid described in the previous report (see abstract to W 86). Since both IV (substance obtained in the present work) and allolupinine described in the previous report are identical, the described acids are identical.
9. CARD:	3/7

PRINTED :
CAT. NO. 1 :
AUG. JOUR. : RAKHIM., No. 22 1959, No. 82139

EDITOR :
PRINT. :
FILE :
ORIG. PUB. :

ABSTRACT
CONT'D : above), 95% oil is obtained which, after chromatography on Al₂O₃, gave IV, m.p. 70-72°,
 η^{20}_D 1.5118; picrate, m.p. 81° (from water),
m.p. 115-123.5° (after drying in vacuum); hydrochloride, m.p. 25.0° (from petroleum-ether
fraction). The other fraction produced a
small quantity of the substance, isomeric IV,
with m.p. below 30°, the structure of which
was not determined. Hydrochloride of IV is

CARD: 5/7

0-39

GROUP#	:	G
CATEGORY	:	
ABG. JOUR.	:	REKOM., No. 23 1950, No. 32230
AUTHOR	:	
INST.	:	
PAGE#	:	
ORIG. PUB.	:	
ABSTRACT cont'd	:	of esters, an ethyl ether, VI (5%), is separated chromatographically; picrate, b.p. 130.5-132°.-- Jan Kovar
CARD:	7/7	

6-10

ERNEST, I.

J. P. (NBS)

Decomposition of diazoketones with cupric oxide. VIII. Preparation of aliphatic β -acryl acrylic acids. I. Ernest and H. Jelinkova (Vysoka skola chem. technol., Prague). *Collection Czechoslov. Chem. Commun.* 24, 3341-7 (1959) (in German); cf. *C.A.* 52, 11806b. —Decompos. of the mixt. of diazoketones RCOCH_2 , Me (I) or Et (II) diazooacetates and CuO powder in C_2H_5 at 70-80° gave alkyl β -acrylates $\text{RCOCH}(\text{CH}_2\text{CO}_2\text{R}')$ (III) (possessing probably the trans configuration as shown by the absorption max. at 982 cm.⁻¹ of the Et ester), (R, R', m. or b.p., % yield given): Pr, Me (IV), m. 37.5°, —, 8; Bu, Et, b.p. 90-3°, 1.4522, 19; Am, Et, b.p. 99.5-100°, 1.4522, 14; $n\text{-C}_8\text{H}_{17}$, Et, b.p. 110-12°, 1.4542, 19; $n\text{-C}_8\text{H}_{17}$, Bu (V), b.p. 127-9°, 1.4537, 16. Me (VI) or Et (VII) fumarates and sym. diketones $\text{RCOCH}(\text{CH}_2\text{CO})\text{R}'$ were obtained as by-products in some cases; if fractionation failed, hydrolysis of the crude mixt. was applied. Heating 30 g. $n\text{-C}_8\text{H}_{17}\text{COCH}_2$, 20.7 g. II, 1000 ml. C_2H_5 , and 2 g. CuO powder with agitation in a water bath at 70° led to a violent decompos.; refluxing then the mixt. 15 min., sepd. the CuO, and evapn. the filtrate in vacuo gave a residue from which was isolated 5.2 g. diacryloylethylene, m. 80.5° (MeOH) and 84.5 g. Liquid; the latter was fractionated to give 6.2 g. VII and 6.2 g. V. Analogous decompos. of 18 g. PrCOCH_2 (VIII) and 18.1 g. II gave a mixt. of VII and $\text{PrCOCH}(\text{CH}_2\text{CO})\text{Et}$ (could not be sepd. by distn.) whereas 21 g. VIII, 14.7 g. I, 200 ml.

Jiri Pantić

C 84

ERNEST, IVAN

1

4-968(AB)

Esters of unsaturated dioxo carboxylic acids. Ivan Erlekt and Zdenka Linhartová. Czech. 88,300, Jan. 10, 1958. Heating a soln. of an alkyl diazomethyl ketone with an α -diazocetic ester in a hydrocarbon solvent in the presence of CuO gives title compds., $\text{RCOCH}:\text{CHCO}(\text{CH}_3)_n\text{CO}_2\text{R}'$ (I), besides $\text{RCOCH}:\text{CHCOR}$ (II) and $\text{RO}_2\text{C}(\text{CH}_3)_n\text{COCH}:\text{CHCO}(\text{CH}_3)_n\text{CO}_2\text{R}$ (III) as side products. I show bacteriostatic and fungicidal activity. Me α -diazolevulin-
ate (IV) from 7.8 g. $\text{MeO}_2\text{C}(\text{CH}_3)_n\text{COCl}$ (cf. C.A. 50, 13749b) heated with stirring with 4 g. MeCOCHN_2 and 3 g. powd. CuO in 1 l. C_6H_6 to boiling, the evolution of N completed by refluxing 15 min., the catalyst filtered off, and the solvent distd. off *in vacuo* gives on standing 1.8-2.2 g. III (R = Me, n = 2), m. 120-1° (MeOH). The liquors fractionated *in vacuo* give 1 g. II (R = Me), b₁ 80-6°, m. 74°, and 2 g. I (R = R' = Me, n = 2), b₁ 119-20° (solidified at -20°). Similarly, IV and Et COCHN_2 give II (R = Et), b₁ 80-6°, m. 52-3° (petr. ether), and 15.6% I (R = Et, n = 2, R' = Me), and Me α -diazoacetylvalerate with Me COCHN_2 gives II (R = Me) and 21% I (R = R' = Me, n = 4), b₁ 130-6°.

L. J. Urbánek

ERNEST, IVAN

Distr: 4E3d
1
Unsaturated 2,5-disubstituted furan derivatives. Ivan
Ernest and Jan Staněk. Czech. 88,700, Feb. 15, 1969.
Treating solns. of $R^1COCH_2CHCO(CH_3)_2R^2$ (I) in an anhyd.

polar solvent with strong mineral acids gives title compds.

$R^1CH_2CH_2CH_2CO_2R^2$ (II), which show bacteriostatic effect. Letting stand a soln. of 2 g. I ($R^1 = CH_3CH_2CO_2Me$, $R^2 = CO_2Me$) in 100 ml. AcOH with 4 drops concd. HCl 24 hrs. at 20° and evapg. the AcOH and HCl at 60° in HCl 24 hrs. at 20° and evapg. the AcOH and HCl at 60° in HCl 24 hrs. at 20° and evapg. the AcOH and HCl at 60° in $R^1 = CH_3CH_2CO_2Me$, $R^2 = CO_2Me$), m. 50° (MeOH). Similarly, 1 g. I ($R^1 = R^2 = CO_2Me$) gives on cooling 1.86 g. II ($R^1 = CH_3CH_2CO_2Me$, $R^2 = CO_2Me$), m. 50° (MeOH). Similarly, 1 g. I ($R^1 = R^2 = Ph$) gives 720 mg. II ($R^1 = CH_3CH_2Ph$, $R^2 = Ph$), m. 60-1° (cyclohexane), and 305 mg. I ($R^1 = R^2 = Ph$) gives 200 mg. II ($R^1 = Me$, $R^2 = CO_2Me$), b. 65-70°, m. 36-7°.

L. J. Urbánek

3
1978 (18)

ERNEST,

Distr: 4E2c(j)/4E3d

✓ Addition of metalloorganic compounds to unsaturated pyridine bases. V. Hněvová and I. Ernest (Vysoká škola chem.-technol., Prague). Collection Czechoslov. Chem. Commun. 25, 1468-74 (1960) (in German).—PhLi, lutidyl-lithium, and EtMgBr added unsatd. 2- and 2,6-substituted pyridine bases having a double bond conjugated with the pyridine nucleus in the side chain. The org. component of the reagent was attached to the β -C atom of the side chain. 2-Methyl-6-styrylpyridine (I) (5 g.) added with stirring to PhLi (from 0.7 g. Li, 5.7 g. PhBr, and 80 ml. Et₂O), the mixt. stirred 1 hr., decompd. with aq. NH₄Cl soln., the basic products extd. with dil. HCl (1:1), basified and extd. with Et₂O gave 3.0 g. 2-methyl-6-(β , β -diphenylethyl)pyridine, m. 177-9°, m. 60.5-7.5° (cyclohexane); picrate m. 184.5-0.5° (EtOH). Similarly, 10 g. I allowed to react with PhLi from 11.5 g. PhBr in 70 ml. Et₂O and 3 g. EtCHO in 100 ml. Et₂O added dropwise gave 4.05 g. 2-methyl-6-(α -benzhydryl- β -hydroxybutyl)pyridine, m. 109.5-70.0° (EtOH); picrate, m. 177-8° (EtOH); O-Ac deriv. m. 110.5-11.5° (EtOH). Analogously the treatment of 2-but enylpyridine (Ia), 2-methyl-6-but enylpyridine (II), 2-styrylpyridine (III), and 2,6-distyrylpyridine with 40-50% excess of 0.5-0.8M PhLi soln. in Et₂O gave, resp., 50.8% 2-(β -pyrido[1,2- α :1',2']pyrazine-6,12-diol (VIII), m. 172-80° (alc.), yield 11.8%. IV and V give VIII. PhMgBr with III in Et₂O-anisole yielded after decompn. with HCl 10.8% α , α -diphenyl-2-piperidinemethanol (IX), m. 81-2° (cf. Tilford, *et al.*, CA 43, 2205g). IX was also obtained by reaction of 5.4 g. Mg and 38.2 g. bromobenzene and addn. of 15.4 g. II in 100 ml. Et₂O in 16.3-h. yield;

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(EtOH-AcOEt)]. The origin of 1,1,3-triphenyl-2,4-bis(1-pyridyl)butane, m. 183-5.5° [dipicrate, m. 138-40° (both from EtOH)], as a side product of V was explained by the reaction of the intermediary organolithium deriv. with another mol. of III. The above addn. reaction failed in the case of 2-vinylpyridine, because of fast polymerization of the unsatd. base, and also in the case of bases having an unsatd. side chain in position 4, e.g. in 4-styrylpyridine, which was recovered unchanged. 2,6-Lutidylolithium yielded with II 49.2% 1,3-bis(0-methyl-2-pyridyl)-2-ethylpropane, b₁ 131.5-5.0°; dipicrate m. 163.5-70.0° (EtOH-Et₂O). In certain cases the above addn. of Li compds. had an analogy in the behavior of Grignard reagents, since I allowed to react with EtBr and Mg in Bu₂O gave IV, whereas reaction of EtMgBr with Ia, II, and III gave complicated mixts. of products. From the reaction mixt. of Ia and EtMgBr was isolated 20.4% 2,4-diethyl-1,3-bis(2-pyridyl)hexane, b₁,₄ 160-70° [dipicrate m. 173-4.5° (Me₂CO)], and from that of II and EtMgBr was obtained 43% 2,4-diethyl-1,3-bis(0-methyl-2-pyridyl)hexane, b₁,₄ 155-70° [dipicrate m. 204-5° (Me₂CO-Et₂O)]. 2-(β -Hydroxybutyl)pyridine, b₁ 117.5°, was obtained in 35.3% yield by adding dropwise a soln. of 27.5 g. EtCHO in 100 ml. Et₂O to the reagent from 49.8 g. 2-picoline and 1 l. 0.6M PhLi soln. in Et₂O, stirring 2 hrs., decompd. with 10% NH₄Cl soln. and working up as usual; metho- ρ -toluenesulfonate m. 121-4° (EtOH-Et₂O).

L. J. Urbášek

COUNTRY : CZECHOSLOVAKIA
CATEGORY : Organic Chemistry. Synthetic Organic Chemistry
ABS. JOUR. : RZKhim., No. 1 1960, No.1138
AUTHOR : Ernest, I.; Šnádel, J.
INST. :
TITLE : Decomposition of Diazoalkanes with Cupric Oxide.
V. A New Reaction of Aliphatic Unsaturated
γ-Diketones
ORIG. PUB. : Collect. Czechosl. Chem. Commun., 1959, 24,
No 2, 530-535
ABSTRACT : No abstract
See RZKhim., No 2, 1960, No 1619.

CARD: 1/1

2-6

ADLEROVA, E.; BLAHA, L.; BOREVICKA, M.; ERNEST, I.; JILEK, J.O.; KAKAC, B.;
NOVAK, L.; RAJSNER, M.; PROTIVA, M.

Synthetic experiments in the group of hypotensive alkaloids. VI.
Some notes on the preparation of alicyclic components in the
synthesis of compounds of the reserpine type. Coll Cz Chem 25 no.1:
221-236 Ja '60. (EEAI 9:2)

1. Forschungsinstitut fur Pharmazie und Biochemie, Prag.
(Alkaloids) (Hypotension)
(Alicyclic compounds) (Reserpine)

HNEVSOVA, V.; ERNEST, I.

lelobine

Synthesis experiments in the series of lelobine alkaloids; synthesis and reactions of 2-styryl-6-butenylpyridine. Coll Cz chem 25 no.3: 748-755 Mr '60. (EEAI 9:12)

1. Institut fur organische Chemie, Technische Hochschule fur Chemie, Prag.

(Alkaloids) (Lelobine) (Butenylstilbazole)

HNEVSOVA, V.; ERNEST, I.

Addition of metallo-organic compounds to unsaturated pyridine bases.
Coll Cz Chem 25 no.5:1468-1474 My '60.

1. Institut fur organische Chemie, Technische Hochschule fur Chemie,
Prag. 2. Jetzige Adresse: Forschungsinstitut fur Pharmazie und
Biochemie, Prag (for Ernest).

NOVAK, L.; JILEK, J. O.; KAKAC, B.; ERNEST, I.; PROTIVA, M.

Synthetic experiments in the group of hypotensive alkaloids. IX. A new method for splitting racemates in the total synthesis of reserpine. Coll Cz Chem 25 no.8:2196-2206 Ag '60. (EEAI 10:9)

1. Forschungsinstitut fur Pharmazie und Biochemie, Prag.

(Alkaloids) (Hypotension) (Tartaric acid)
(Reserpine)

ADLEROVA, E.; ERNEST, I.; HNEVSOVA, V.; JILEK, J.O.; NOVAK, L.; POMYKECEK, J.; RAJSNER, M.; SOVA, J.; VEJDELEK, Z.J.; PROTIVA, M.

Experiments on synthesis in the group of hypotensive alkaloids.
VIII. Syntheses of some tryptamine derivatives, substituted in
positions 5,6, and 7. Coll Cz chem 25 no.3:784-796 Mr '60.
(EEAI 9:12)

1. Forschungsinstitut fur Pharmazie und Biochemie, Prag.
(Alkaloids) (Aminoethylindole) (Hypotension)

IOTA, C.G.; RUNCAN, V.; CHITESCU, Elena; SUTEANU, St.; ERNEST, I.

The neuov vegetative syndrome in chronic hepatitis. I. Preliminary investigations. Stud. cercet. med. intern. 2 no.2:203-217 '61.
(HEPATITIS, INFECTIOUS complications)
(AUTONOMIC NERVOUS SYSTEM diseases)

JILEK, J. O.; ERNEST, I.; NOVAK, L.; RAJSNER, M.; PROTIVA, M.

Synthetic experiments in the group of hypotensive action alkaloids.
XII. Contribution to the terminal phases of total synthesis of
reserpine and deserpidine. Coll Cz Chem 26 no.3:687-700 Mr '61.
(EEAI 10:9)

1. Forschungsinstitut fur Pharmazie und Biochemie, Prag.

(Reserpine) (Deserpidine) (Alkaloids)

SKODA, J.; ERNEST, I.; STANEK, J.; HABERMANN, V.

The relationship between structure and antibacterial effect of α -unsaturated γ -diketones. Coll Cz Chem 26 no.3:874-880 Mr '61.
(EEAI 10:9)

1. Institute of Organic Chemistry and Biochemistry, Czechoslovak Academy of Science, Prague, Department of Organic Chemistry, Institute of Chemical Technology, Prague, and the Institute for Clinical Chemistry, Medical Faculty of the Charles University, Plzen. 2. Present Address: Research Institute for Pharmacy and Biochemistry, Prague (for Ernest)

(Ketones) (Bactericidal action) (Unsaturated compounds)

ERNEST, I.; STANEK, J.

Decomposition of diazoketone with copper (II) oxide, Part 9: Kinetics of cyclization of unsaturated γ -diketone. Coll Cz Chem 26 no.4: 1039-1047 Ap '61.

1. Institut fur organische Chemie, Technische Hochschule fur Chemie, Prag (for Stanek) 2. Forschungsinstitut fur Pharmazie und Biochemie, Prag (for Ernest)

(Diazocompounds) (Copper oxides) (Ketones)

ERNEST, I.; PROTIVA, M.

Synthetic tests in the group of hypotensive active alkaloids. Part
14: (+)-methyl-O-(α -carbethoxysyringoyl)-10-methoxydeserpidat.
Coll Cz Chem 26 no.4:1137-1144 Ap '61.

1. Forschungsinstitut fur Pharmazie und Biochemie, Prag.

(Alkaloids)

ERNEST, I.

"Electronic Theories of Organic Chemistry" by J. W. Baker. Reviewed
by I. Ernest; Coll. Gen. Chem. 26 no. 4: 1215-1216 Ap '61.

(Baker, J. W.) (Chemistry, Organic)
(Electronics)

ERNEST, I.; JILEK, J.O.; VEJDELEK, Z.J.; PROTIVA, M.

Sythetic experiments in the group of active hypotensive alkaloids.
Pt. 26. Coll Cz Chem 28 no.4:1022-1030 Ap '63.

1. Forschungsinstitut fur Pharmazie und Biochemie, Prag.

ERNEST, I.

Rules of carotenoid nomenclature. Chem listy 57 no.4:348-349
Ap '63.

ERNST, J.; HEBKY, J.

Discussion on teaching chemistry in secondary and higher schools.
Chem Listy 58 no. 9:1129-1130 S '64.

ERNST, I.; KEST, B.

Synthetic tests on the group of blood pressure reducing alkaloids. Pt.34. Chem & Chem 29 no.11:2663-2680 N '64.

I. Forschungsinstitut fur Pharmazie und Biochemie, Prague.

PROTIVA, M., inz. dr. DrSc. (Praha 3, Kourimska 17); NOVAK, L.;
VEJDELEK, Z.J.; ERNEST, I.

Sympathetic ganglionic blocking agents. Pt.14. Česk. farm.
14 no.7:346-351 S '65.

ERNEST, I.

"Organic chemistry" by R.Lukes. Reviewed by I.Ernest. Coll
Cz Chem 30 no.3:920 Mr '65.

ERNEST, I.; HEBKY, J.

Discussion on the work with small quantities in organic chemistry.
Chem listy 59 no.5:633-634 My '65.

CZECHOSLOVAKIA

ERNEST, I; KAKAC, B.

Research Institute for Pharmacy and Biochemistry (Forschungs-
institut fur Pharmazie und Biochemie), Prague

Prague, Collection of Czechoslovak Chemical Communications,
No 1, January 1966, pp 279-290

"Experiment with synthesis in the group of hypotensive active
alkaloids. Part 38: Synthesis of the \pm apiochimbine and some
additional iohimbane derivatives."

JIRKOVSKY, I.; PROTIVA, M.; ERNEST, J.

Synthetic experiments in the group of active hypotensive alkaloids. Pts. 29-30. Coll Cz Chem 28 no. 11: 3096-3112 N° 63.

1. Forschungsinstitut fur Pharmazie and Biochemie, Prag.